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Non-precious Co₃O₄-TiO₂/Ti cathode based electrocatalytic nitrate reduction: Preparation, performance and mechanism



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ABSTRACT

The presence of high nitrate (NO₃⁻) concentration in natural water constitutes a serious issue to the environment and human health. Therefore, the development of low-cost, stable non-precious metal catalysts is imminent for efficient NO₃- reduction. In this study, we prepared a Co₃O₄-TiO₂/Ti cathode via combining sol-gel and calcination methods and evaluated its performance for electrocatalytic NO₃- reduction. The dispersion of the Co₃O₄ catalyst particles was improved by the addition of PVP to the coating liquid. The presence of anatase could effectively stabilize Co_3O_4 and prevent the releasing of toxic Co ions into the solution. The Co_3O_4 - TiO_2/Ti cathode with the optimized performance for NO₃- reduction could be prepared by four times coating at calcination temperature of 500 °C. The electrocatalytic reduction of NO₃- was negligibly impacted by solution pH in the range of 3.0-9.0, while it could be facilitated by elevating the current density from 2.5 to 25 mA cm². Ammonium ions were the main final NO₃- reduction product, and the presence of Cl- was capable to oxidize ammonium ions to N2 due to the electrochemical production of reactive chlorine species. The electrochemical analyses, scavenging experiments and density functional theory calculations collectively confirm that NO₃- reduction was mainly induced by the Co²⁺-Co³⁺-Co²⁺ redox process instead of being directly resulted from the electrons generated at the cathode. Unlike noble metal (e.g., Pd and Ag) based catalytic reaction systems, in the present Co₃O₄ mediated electrocatalytic reaction process, atomic H* would more favorably turn to H₂ by Heyrovsky and Tafel routes and therefore contributed marginally to the NO3- reduction. Generally, this study provided a new paradigm for designing the stable and cost-effective cathode for NO₃- reduction.

1. Introduction

In water, nitrate (NO_3^-) is considered an underlying hazardous material that is derived from various sources including the feedlot runoff, the excessive use of nitrogen fertilizer and natural degradation of pollutants [1]. The presence of NO_3^- in drinking water can cause serious health problems such as the cancers, liver damage and blue baby syndrome due to its transformation product, i.e. nitrite anions (NO_2^-) [2]. To diminish this potential health risk, the concentration limit for NO_3^- in drinking water is set at 10 and 11.3 ppm by the United States EPA and the European Drinking Water Directive, respectively [3,4]. In this regard, the removal technologies of NO_3^- are imperious demands in populated regions.

Physicochemical treatments such as electrodialysis, ion exchange

and reverse osmosis are the most common technologies for NO_3^- removal from water [5], but the post-treatments are still needed to address the concentrated solutions produced in these methods. Although biological denitrification can gradually convert NO_3^- to N_2 [6], it is a time-consuming process and its performance is strongly impacted by the variable conditions, such as dissolved oxygen and temperature. Recently, zero valent metals (e.g. iron [7], aluminum [8], and magnesium [9]) based stoichiometrically chemical reduction of NO_3^- has been increasingly explored. Unfortunately, the flexibility of this strategy is greatly confined by the formation of inert oxide surface, deteriorating the electron transfer efficiency, and the inevitable production of secondary solid sludge. More recently, some catalytic hydrogenation processes with the combined noble/promoting metals (e.g., Pd/Cu and Pd/Fe) have been proposed for NO_3^- reduction

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[10,11]. In these processes, the reduction of NO_3^- is mainly initiated by promoter metal (Eq. (1)) with the formation of NO_2^- , and the subsequent reduction process is gradually achieved by the noble metal activated hydrogen (Eq. (2)) [12]. However, catalytic hydrogenation reduction requires H_2 as a reducing agent or electron donor, and there are safety hazards in the transport and use of H_2 , which greatly limits its industrial application potential [13].

$$NO_3^- + 2M[] \rightarrow [M_2O]_{surf} + NO_2^-$$
 (1)

$$H_2 + 2Pd[] \rightarrow 2Pd[H] \tag{2}$$

Considering the advantages of self-produced hydrogen, environmental compatibility and ambient operating conditions, electrocatalysis has been regarded as a promising method for denitrification [14], in which the reduction of NO₃ can be obtained by either a direct or an indirect pathway at the cathode surface. Direct reduction of NO₃⁻ is initiated by the electron tunneling or generation of chemisorbed complex between the NO₃⁻ and cathode whereas in the indirect pathway, NO₃ is reduced by a surface-adsorbed atom H*, which is produced from the reduction of protons by electron via the Volmer process [15]. Specially, Pd-based catalysts have recently been distinguished mainly because of their potential to electrochemically reduce oxyanion contaminants in water in which the indirect reduction mediated by atomic H* plays a leading role compared to the direct electron transfer from the cathode to NO₃ [16]. Nevertheless, considering the extremely expensive price of Pd, the research significance of the Pd-based catalyst is obviously theoretical but not practical. Additionally, despite the fact that the indirect reduction mechanism of NO₃⁻ has been well explored in electrochemical noble metal-based process, the targeted strategies to develop some specific catalysts for the direct reduction of NO₃ is

The cost-effective cobalt oxide (Co₃O₄) is a typical p-type metal oxide, which exhibits a normal spinel structure with Co²⁺ ions in tetrahedral interstices and Co³⁺ ions in octahedral interstices. During the recent decades, Co₃O₄ has been demonstrated to own strong catalytic reduction activity and thus extensively used in electrochemical hydrogen evolution reaction [17] and reduction of CO2 [18] and NOx [19]. Recently, Su et al. [20] reported that the pure Co₃O₄ loaded onto the titanium substrate by thermal sintering could also electrocatalytically reduce NO₃ in the water. However, the Co₃O₄-mediated electrochemical mechanism of NO₃ reduction has remained unclear. Moreover, it is known that the biologically toxic cobalt ion probably causes human carcinogen, and the discharge standard of cobalt ions in industrial wastewater is 1 mg L⁻¹ based on GB/T 25467-2010 of China. Therefore, efforts should be made to improve the stability of the cobaltbased catalyst and prevent its leaching from the substrate and leaking into the water. To this end, we prepared a novel cathode, Co₃O₄-TiO₂/ Ti, by combining sol-gel and calcination methods, which seems to be promising for remarkably facilitate the electrocatalytic reduction of NO₃. The effects of electrode preparation factors (calcination temperature and coating times) and operation parameters (pH, current density and chloride ions concentration) on NO₃⁻ reduction were also investigated. In addition, based on the characterization of the prepared catalysts using XRD, SEM, HRTEM, EDS elemental mapping and XPS, we tried to relate the results to the catalytic activity. Furthermore, electrochemical analyses, radical scavenging experiments and density functional theory calculations were utilized to inspect the possible NO₃ reduction mechanisms.

2. Experimental section

2.1. Reagents and materials

Titanium (Ti) plate (99.99% purity, thickness $1.0\,\mathrm{mm}$) and IrO_2 -RuO $_2$ /Ti plate were purchased from Yunxuan Metallic Materials Co. Ltd. Potassium iodide (KI, > 99%), mercuric iodide red

 $(HgI_2, > 99.5\%),$ potassium sodium tartrate tetrahydrate $(NaKC_4H_4O_6:4H_2O_7) > 99\%$, sulfamic acid $(NH_2SO_3H_7) > 99\%$, sulfanilamide ($C_6H_8N_2O_2S$, > 99%), titanium butoxide ($C_{16}H_{36}O_4Ti$), cobalt hexahydrate $(Co(NO_3)_2 \cdot 6H_2O, > 99\%),$ sodium chloride (NaCl, > 99.5%), sodium pyrophosphate (Na₄P₂O₇, > 99%), cobaltic oxide (Co₂O₃, > 99%), sodium acetate (C₂H₃NaO₂, > 99%), ethanol (95%), hexadecyl trimethyl ammonium bromide (C₁₉H₄₂BrN, TBA, > 99%), polyvinyl pyrrolidone K30 ((C₆H₉NO)_n, PVP), ammonium chloride (NH₄Cl, 99.8%), potassium nitrate (KNO₃, > 99.5%), sodium nitrite (NaNO₃, > 99%), H₂SO₄ (95%–98%), HCl (36%–38%), NaOH (> 96%) were provided by Sinopharm Chemical Reagent Co. Ltd., China, All the reagents were analytical grade and used without further purification. All working solutions were prepared using ultrapure water (18.2 M Ω cm resistivity). The experiments were done triplicate.

2.2. Preparation of the working cathode

The substrate of electrode was Ti plate with size of 2.5×5.0 cm. The Ti plate was first polished with 300-mesh emery paper and then boiled in 20% sulfuric acid for 40 min. Next, the Ti plate was cleaned by ultrapure water, sonicated for 15 min, and dried for use. To synthesize the typical Co₃O₄-TiO₂/Ti cathode, 0.5 mL titanium butoxide, 0.1 mL hydrochloric acid and $0.15\,\text{mL}$ ultrapure water were dissolved in $8.5\,\text{mL}$ ethyl alcohol. Then, 0.5 g PVP was added and stirred vigorously for 0.5 h. Subsequently, 2.69 g Co(NO₃)₂•6H₂O (1 M) was added and stirred to obtain a purplish red liquid. The prepared sol-gel was pasted to the both faces of the pretreated Ti plates using a brush. The painted Ti plate was dried at 100 °C for 10 min. Then it was sintered at 500 °C for 2 h in a muffle oven to remove all organic substances with the formation of the oxide. The above procedures were repeated 4 times and the electrode films were annealed for 6 h at 500 °C in the last time. Other different kinds of cathodes were prepared with the same process with that for the Co₃O₄-TiO₂/Ti (without PVP) cathode. Furthermore, the coating times (2, 4, 6 and 8) and the calcination temperatures (300 °C, 400 °C, 500 °C, 600 °C and 700 °C) were adjusted to optimize the preparation

2.3. Electrocatalytic reduction of NO₃

A DC potentiostat (GPS-3030D, Gwinstek, China) was utilized as the power supply for electrochemical experiments. The electrocatalytic NO $_3^-$ reduction experiments were carried out in a 150 mL electrolytic cell at room temperature (25 °C). If not specially mentioned, the volume of the working solution was 100 mL, NO $_3^-$ concentration was 50 mg L 1 , and 0.1 M Na $_2$ SO $_4$ was used as supporting electrolyte to maintain the conductivity of the solution. The solution pH was adjusted using concentrated NaOH and H $_2$ SO $_4$. The 12.5 cm 2 Co $_3$ O $_4$ -TiO $_2$ -PVP electrodes were used as cathode and 12.5 cm 2 IrO $_2$ -RuO $_2$ /Ti electrodes were used as anode. The distance between the electrodes was 2 cm. The typical electrochemical experiment was conducted at current density of 10 mA cm 2 . All experiments were repeated at least twice.

2.4. Physicochemical characterizations and DFT calculation

The crystalline nature of the surface of ${\rm Co_3O_4\text{-}TiO_2/Ti}$ cathode was examined by X-ray diffraction (XRD) measurement and the 20 values acquired within the range of 10° to 80° using a Bruker D2 PHASER X-ray diffractometer that uses Cu ${\rm K}_\alpha$ radiation. The morphology of the cathode surface was depicted by a scanning electron microscopy (SEM, FEI QUANTA FEG250) and high-resolution transmission electron microscope (HRTEM, JEM-2100 F). The X-ray photoelectron spectroscopy (XPS) was obtained using a Thermo Fisher ESCALAB 250Xi system with Al ${\rm K}_\alpha$ as an X-ray source. C 1s peak at 284.8 eV from residual carbon was conducted to correct the binding energy of other elements. The details of DFT calculation were provided in the supplementary material.

2.5. Electrochemical characterizations

All electrochemical measurements were performed on an electrochemical station (PGSTAT 302 N, Metrohm). A standard three-electrode system was used in which the working electrode was a home made $\rm Co_3O_4\text{-}TiO_2\text{-}/Ti$ cathode, the counter electrode was a platinum plate, and the reference electrode was a saturated calomel electrode (SCE). The effective working area of the working and counter electrodes was $12.5~\rm cm^2$. The electrolyte for linear sweep voltammetry (LSV) and cyclic voltammetry (CV) tests was composed of $0.1~\rm M~Na_2SO_4$ solution with or without $50~\rm mg~L^{-1}~NO_3$ -. CV tests were recorded between $0.0~\rm V$ and $-1.5~\rm V/SCE$ at a scan rate of $20~\rm mV~s^{-1}$. Electrochemical impedance spectroscopy (EIS) measurements were operated in $0.1~\rm M~Na_2SO_4$ solution using the frequency ranged from $10^5~\rm to~10^{-2}~Hz$ and the amplitude of the potential of $10~\rm mV$.

2.6. Chemical analyses

The concentrations of NO₃⁻, NO₂⁻ and NH₄⁺ were determined with a UV–vis spectrophotometer as it has been depicted in detail in the previous literature [21]. The residual NO₃⁻ ratio $(R_{NO_3}^-)$, the generated NO₂⁻ ratio $(S_{NO_2}^-)$, the generated NH₄⁺ ratio, $(S_{NO_4}^+)$, the total nitrogen (TN) removal efficiency, (R_{TN}) and N₂ selectivity $(S_{N_2}$, were calculated by the following equations:

$$R_{NO_3^-}(\%) = \frac{[NO_3^-]_t}{[NO_3^-]_0} \times 100$$
 (3)

$$S_{NO_2^-}(\%) = \frac{[NO_2^-]_t}{[NO_3^-]_0} \times 100$$
 (4)

$$S_{NH_4^+}(\%) = \frac{[NH_4^+]_t}{[NO_3^-]_0} \times 100 \tag{5}$$

$$R_{TN} = 1 - R_{NO_3^-} - S_{NO_2^-} - S_{NH_4^+}$$
 (6)

$$S_{N_2}(\%) = \frac{R_{TN}}{1 - R_{NO_3^-}} \times 100 \tag{7}$$

where $[NO_3^-]_0$ is the initial concentration of NO_3^- and $[NO_3^-]_t$, $[NO_2^-]_t$ and $[NH_4^+]_t$ are the concentrations of NO_3^- , NO_2^- and NH_4^+ at time t.

3. Results and discussion

3.1. Catalytic NO₃⁻ reduction by different cathodes

The comparative NO₃ reduction process efficiency at different cathodes was depicted in Fig. 1. As shown in Fig. 1(a), the NO₃ removal efficiency of ${\rm TiO_2/Ti}$ cathode was only 45% at current density of 10 mA cm⁻² after 120 min. However, as for Co₃O₄-TiO₂/Ti (without PVP) cathode, NO₃ removal efficiency suddenly became twice that of TiO₂/Ti cathode, reaching approximately 80%. Moreover, with the introduction of the dispersant PVP, the removal efficiency of NO₃ was further increased to 89%. Although the addition of TiO2 resulted in a slight decrease (nearly 6%) in the removal efficiency of NO₃⁻ compared to Co₃O₄/Ti cathode, the release of Co ions into the aqueous solution was remarkably inhibited, probably due to the stabilization effect of TiO2 (see Fig. 1(b)). In the case of Co3O4/Ti cathode, the leaching of Co ions (Co²⁺/Co³⁺) reached 0.2 mg L⁻¹ after 120 min, while Co ions were barely detected for both the Co₃O₄-TiO₂/Ti and Co₃O₄-TiO₂/Ti (without PVP) cathodes. Recently, many non-precious catalysts for NO₃ reduction have been developed as the alternatives to conventional noble Pd catalyst. As shown in Table S1, the Co₃O₄-TiO₂/ Ti cathode exhibits significantly superior NO₃ reduction activity over other non-precious electrodes.

As shown in Fig. 1(c), the profiles of ammonium ($\mathrm{NH_4}^+$) formation as a function of electrolysis time have similar trends with those of $\mathrm{NO_3}^-$

removal in the above reaction systems. The amount of NH_4^+ formed in the case of Co_3O_4 /Ti cathode was larger than those of Co_3O_4 -TiO₂/Ti and Co_3O_4 -TiO₂/Ti (without PVP) cathodes. In addition, among all the cathodes tested, the TiO_2 /Ti produced the lowest amount of NH_4^+ , due to its inferior performance for NO_3^- reduction. During NO_3^- reduction process, the toxic intermediate NO_2^- was negligibly detected in the cases of Co_3O_4 -TiO₂/Ti, Co_3O_4 -TiO₂/Ti (without PVP) and Co_3O_4 /Ti cathodes, while a significant amount (around 1%) of NO_3^- was transformed into NO_2^- when using TiO_2 /Ti as the cathode. Although NO_3^- reduction efficiency of Co_3O_4 -TiO₂/Ti cathode was slightly lower than that of Co_3O_4 /Ti cathode, the TN removal efficiency and N_2 selectivity for the former were comparable to that of the latter as shown in Fig. S1. Besides, the TN removal efficiency and N_2 selectivity of Co_3O_4 -TiO₂/Ti (without PVP) and TiO_2 /Ti cathodes were much lower than those of Co_3O_4 -TiO₂/Ti cathode.

Fig. 2 shows the XRD pattern of Co_3O_4 - TiO_2 /Ti cathode calcinated at the temperature of 500 °C. A comparison was performed between the obtained XRD data and standard patterns of various cobalt oxide and titanium oxide structures including cubic phase Co_3O_4 (PDF#42-1467), Ti (PDF#44-1294), anatase TiO_2 (PDF#21-1272), and rutile TiO_2 (PDF#21-1276). It is observed that all the detectable peaks agree well with standard cubic phase Co_3O_4 , anatase TiO_2 , and rutile TiO_2 . And the main characteristic peaks of Co_3O_4 are at $2\theta = 31.3^\circ$, 36.9° , 38.6° , 44.8° , 59.4° , 65.2° , 77.3° , which correspond to (220), (311), (222), (400), (511), (440), and (533) planes, respectively. This shows that the main components of the as-prepared cathode were the target catalyst Co_3O_4 and the dispersant carrier TiO_2 .

SEM images (Fig. 3) illustrate the morphology of as-prepared catalysts. Fig. 3(a)-(c) correspond to Co₃O₄-TiO₂/Ti, Co₃O₄-TiO₂/Ti (without PVP), and Co₃O₄/Ti cathodes, respectively. It can be clearly seen that Co₃O₄ particles on Co₃O₄-TiO₂/Ti cathode were small and uniform, whereas that on Co₃O₄-TiO₂/Ti (without PVP) and Co₃O₄/Ti cathodes were apparently agglomerated and represented in the forms of disordered nanotubes and irregular aggregates, respectively, showing the successful role of PVP as surface dispersant by mixing PVP and metal-salt precursor solution [22]. Fig. 3(d)-(h) provide the EDS maps of Co and Ti elements at the above three cathodes. Obviously, the presence of PVP could effectively improve the uniformity of Co and Ti distribution on the surface of Co₃O₄-TiO₂/Ti cathode, which was therefore better than that of Co₃O₄-TiO₂/Ti (without PVP) cathode. As for Co₃O₄/Ti cathode, the aggregation of Co element was more pronounced in comparison with the other two cathodes, indicating the worst distribution of Co₃O₄ on Co₃O₄/Ti cathode. The electrocatalytic property could be improved by using PVP as surface stabilizer and growth modifier since it could effectively alleviate the agglomeration of the catalysts and enhance the relative population of stable clusters during the synthesis of nanoparticles (NPs), thereby more active sites are exposed at the cathode surface. [23,24] Therefore, it can be concluded that both the TiO2 and the PVP dispersant could evidently promote the distribution and stability of Co₃O₄ on the Ti plate, which will expose more active sites and inhibit the leaching of catalyst from the electrode surface. This may explain the superior catalytic performance and stability of Co₃O₄-TiO₂/Ti cathode over Co₃O₄-TiO₂/Ti (without PVP) and Co₃O₄/Ti cathodes as shown in the Fig. 1. HRTEM images of Co₃O₄-TiO₂/Ti in Fig. 4(a) presents dark Co₃O₄ species with a 10-50 nm diameter. The primarily exposed crystal planes of as-synthesized Co₃O₄ are (311) with the lattice spacing (d) of 0.244 nm (Fig. 4(b)), which is consistent with the XRD patterns.

3.2. Preparation parameters

3.2.1. Effect of coating times

The Co ions concentration of 0, 0.5, 1.0, and 2.0 M with 4 coating times was investigated. The NO_3^- concentration decay kinetics and LSV curves (Fig. S2) demonstrate that 1.0 M was compatible. In this section, the studied number of coating times was 2, 4, 6, and 8 with Co

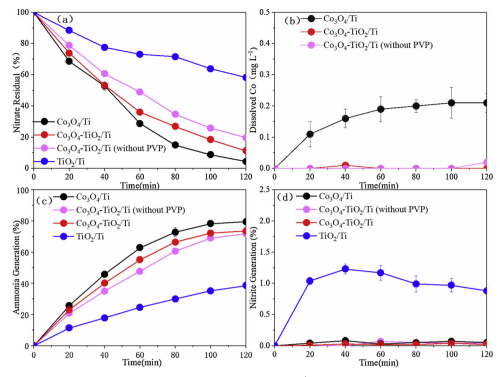


Fig. 1. NO_3^- concentration decay kinetics (a), and evolution of concentration of Co (b), NH_4^+ (c) and NO_2^- (d) during electrolysis time. Experimental conditions: $[NO_3^-]_0 = 50 \text{ mg L}^{-1}$, pH = 7.0, current density = 10 mA cm⁻².

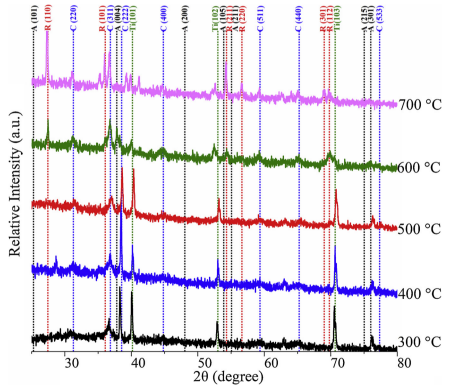


Fig. 2. XRD patterns of Co_3O_4 -TiO₂/Ti cathodes calcined at 300 °C, 400 °C, 500 °C, 600 °C and 700 °C. The locations of the characteristic peaks for Co_3O_4 (C), Ti substrate (Ti), anatase (A), and rutile (R) are represented by the vertical dashed lines.

ions concentration of 1.0 M, which was represented as 2-Co₃O₄-TiO₂/Ti, 4-Co₃O₄-TiO₂/Ti, 6-Co₃O₄-TiO₂/Ti and 8-Co₃O₄-TiO₂/Ti cathodes, respectively. Although, as shown in Fig. S3, the coating times negligibly affected the morphology of the electrode surface, too many or too few coating times were not conducive to $\mathrm{NO_3}^-$ reduction. The $\mathrm{NO_3}^-$

removal efficiencies for $2\text{-Co}_3\text{O}_4\text{-TiO}_2/\text{Ti}$, $4\text{-Co}_3\text{O}_4\text{-TiO}_2/\text{Ti}$, $6\text{-Co}_3\text{O}_4\text{-TiO}_2/\text{Ti}$ and $8\text{-Co}_3\text{O}_4\text{-TiO}_2/\text{Ti}$ cathodes were 80%, 89%, 64% and 61%, respectively, within 120 min. Thus, the optimum number of coating times was 4. As illustrated in Fig. 5(b), $4\text{-Co}_3\text{O}_4\text{-TiO}_2/\text{Ti}$ cathode has the largest response current intensity. And the trend of response current

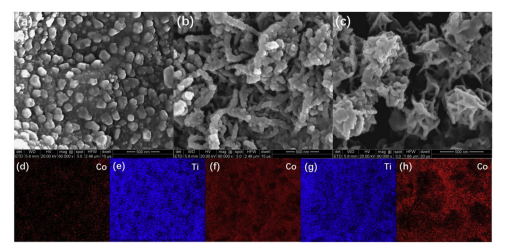


Fig. 3. SEM images of Co_3O_4 -Ti O_2 /Ti (a), (b) Co_3O_4 -Ti O_2 /Ti (without PVP) (b) and Co_3O_4 /Ti (c) cathodes. Elemental mapping of Co_3O_4 -Ti O_2 /Ti (without PVP) (f,g) Co_3O_4 /Ti (h) cathodes. Elemental mapping: (d-h) correspond to Co, Ti, Co, Ti and Co maps, respectively.

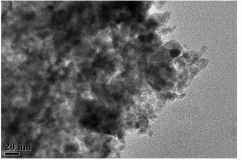
intensity at the same voltage was consistent with the NO_3 reduction trend of Fig. 5(a). The above experimental and LSV results can be explained by the fact that when the number of coating times was below 4, insufficient Co loadings may not afford enough active sites for NO_3 reduction. In contrast, when the number of coating times was more than 4, the catalyst coating on the Ti plate was thick, which probably deteriorated the electron transfer, and the increased potential would exacerbate the occurrence of hydrogen evolution side reactions, therefore led to a decrease in NO_3 reduction efficiency.

3.2.2. Effect of calcination temperature

In this section, the effect of calcination temperatures on the performance of Co₃O₄-TiO₂/Ti cathode was investigated. The as-prepared cathodes sintered at 300 °C, 400 °C, 500 °C, 600 °C and 700 °C were marked as 300-Co₃O₄-TiO₂/Ti, 400-Co₃O₄-TiO₂/Ti, 500-Co₃O₄-TiO₂/ Ti, 600-Co₃O₄-TiO₂/Ti and 700-Co₃O₄-TiO₂/Ti, respectively. Fig. 6(a) shows that there were significant differences in NO₃ reduction efficiency between the cathodes sintered at different temperatures. The optimum NO3- removal efficiency was observed for the cathode calcination in the temperature range of 300-500 °C, e.g., approximately 89% for 500-Co₃O₄-TiO₂/Ti cathode. A moderate NO₃ removal efficiency was obtained for 600-Co₃O₄-TiO₂/Ti cathode (67%) while the 700-Co₃O₄-TiO₂/Ti cathode had the marginal NO₃ removal efficiency, approximately 34%. The variation trend of response current intensity observed from the LSV measurements (Fig. 6(b)) was in good accordance with that of the cathodes for NO₃ removal, which confirmed the best performance of the 500-Co₃O₄-TiO₂-PVP cathode.

As shown in Fig. 2, the main characteristic peaks of Co_3O_4 in XRD spectra for $\text{Co}_3\text{O}_4\text{-TiO}_2/\text{Ti}$ cathode became narrower and sharper as the calcination temperature increased in the range of 300–700 °C. According to the previous studies, complete conversion of Co_3O_4 can be

achieved in the sol-gel and calcination process when the temperature exceeds 270 °C [20]. Better crystallinity structure of Co₃O₄ could be obtained at higher calcination temperatures. Nevertheless, Ti plate has some problems when being sintered at relatively high temperature. Specially, as demonstrated in the previous literature [20], the conductivity of Ti plate was greatly reduced when the calcination temperature reached 600 °C. Besides, the phase structure, composition, and crystallinity of TiO2 are of great influence on its electrocatalytic activity and electrochemical properties [25]. Therefore, XRD was also used to analyze the changes of TiO2 phase on the cathodes sintered at various temperatures. It can be seen from the XRD patterns presented in Fig. 2 that the calcination temperature significantly impacted the crystallization and phase structures of surface catalysts on Co₃O₄-TiO₂/Ti cathode. For Co₃O₄-TiO₂/Ti cathodes sintered at the temperature of 300-500 °C, except the diffraction peaks of Ti substrate and Co₃O₄, two broad peaks at $2\theta = 53.9^{\circ}$ and 76.0° can be attributed to the (105) and (301) plane diffraction, respectively, of anatase TiO2. However, as for $600\text{-Co}_3\text{O}_4\text{-TiO}_2/\text{Ti}$ cathode, small peaks at $2\theta = 27.4^\circ$, 36.1° , 54.3° , 56.6°, 69.0°, and 69.8° correspond to (110), (101), (211), (220), (301), and (112) plane diffraction, respectively, of rutile. Notably, the intensity of rutile diffraction peaks of Co₃O₄-TiO₂/Ti cathode increases sharply at the calcination temperature of 700 °C. This suggests the occurrence of phase transformation of anatase to rutile when elevating the calcination temperature from 300 to 600 °C, which was even more pronounced for 700 °C. Furthermore, with the calcination temperature increase to 600 °C and 700 °C, the crystal turned to aggregation, which was probably responsible for the decrease of NO₃ reduction efficiency (Fig. S4). Based on these results, it can be concluded that the phase transformation of TiO2 began to take place at the calcination temperature above 500 °C. When calcination temperature reaches 600 °C, the rutile phase even became the prevalent phase of the composite. This



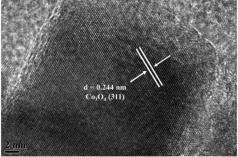


Fig. 4. HRTEM images of Co₃O₄-TiO₂/Ti.

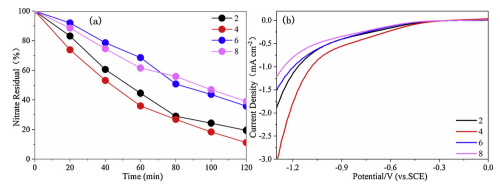


Fig. 5. NO_3^- concentration decay kinetics (a) and LSV curves (b) for different coating times. Experimental conditions: (a) $[NO_3^-]_0 = 50 \text{ mg L}^{-1}$, pH = 7.0, current density = 10 mA cm⁻², (b) 0.1 M $SO_4^- + 50 \text{ mg L}^{-1}$ NO_3^- solution, scan rate: 100 mV s⁻¹).

was consistent with the results in the previous literature papers [26].

Based on the EIS measurements, the Nyquist plots of various Co₃O₄-TiO₂/Ti cathodes sintered at various temperatures are provided in Fig. 6(c). The radius of semicircle was related with the charge transfer resistance and the sample with smaller arc radius indicates faster interfacial electron transfer [27]. The calcination temperature exerts a significant effect on the electrochemical properties of the electrodes. In this study, when the calcination temperature was increased from 300 to 500 °C, the radius of the arc did not appear, indicating that the charge transfer resistance was small. However, when the calcination temperature rose to 600 °C, the arc radius appeared, suggesting the fact that increasing calcination temperature caused an increase in impedance for Co₃O₄-TiO₂/Ti cathode. And the charge transfer resistance of Co₃O₄-TiO₂/Ti cathode obtained at the calcination temperature of 700 °C was larger than that of 600 °C. The change of EIS results indicates that the phase transformation of anatase to rutile with elevating calcination temperature could inhibit the charge transfer within the cathode, which adverse to NO₃ reduction [28]. This can be ascribed to the higher electric resistance of rutile than that of anatase [28].

Therefore, the relatively low NO_3^- reduction efficiencies for 600-Co₃O₄-TiO₂/Ti and 700-Co₃O₄-TiO₂/Ti cathodes could be explained by the increase in the charge transfer resistance of Co_3O_4 -TiO₂/Ti cathode prepared at a high calcination temperature. Consequently, appreciable Co_3O_4 -TiO₂/Ti cathodes could be obtained at the calcination temperature of 300–500 °C, in which cobalt precursor could be completely transformed to Co_3O_4 and the relatively low charge transfer resistance could be also ensured.

3.3. Effect of electrochemical reaction parameters

3.3.1. Effect of solution pH and current density

As illustrated in Fig. 7, the effect of solution pH (3.0–9.0) on NO_3^- reduction was evaluated in this section. Fig. 7(a) and (b) show the reduction curves of NO_3^- over time and the final solution pH for different initial pH values. Contrarily to the previous studies [29], Fig. 7(a) shows that different pH conditions did not play significant role on NO_3^- reduction efficiency over the electrolysis time. This indicates that Co_3O_4 -TiO₂/Ti cathode prepared in this study has a wide working pH range and can be potentially used to treat the NO_3^- containing wastewaters in a relatively large pH range. Besides, Fig. 7(b) shows that, regardless of the initial pH, the pH value was significantly increased after 120 min reaction. For example, after 2 h of nitrate reduction, the solution pH increased from 3.0, 5.0, 7.0 and 9.0 to 9.0, 10.6, 10.9 and 11.0, respectively. The explanation for this pH variation will be discussed in the following Section 3.4.

On the other hand, the reduction efficiency of NO_3^- was significantly enhanced with the increase of current density from 2.5 to $10~\text{mA}~\text{cm}^{-2}$ (Fig. 8(a)). However, when the current density was increased from $10~\text{to}~25~\text{mA}~\text{cm}^{-2}$, the removal efficiency of NO_3^- was not

greatly improved. Fig. S5 shows that the reduction of $50\,\mathrm{mg\,L^{-1}\,NO_3}^-$ followed a zero-order kinetic trend at the current density of 2.5 and $5\,\mathrm{mA\,cm^{-2}}$ with the rate constant increased from 0.190 (R² = 0.99) to 0.440 (R² = 0.98) min (indicating a charge controlled kinetic stage) while a first-order kinetic trend was observed at the current densities of 10 and 25 mA cm² (indicating a mass controlled kinetic stage) with the rate constant of 0.018 (R² = 0.99) and 0.027 (R² = 0.99) min , respectively. As the current density was increased from 10 to 25 mA cm², more energy was used to generate hydrogen bubble, so that the increase in the NO₃ removal efficiency was not significant as expected. There is a good correspondence between the amount of $\mathrm{NH_4}^+$ produced and the amount of $\mathrm{NO_3}^-$ reduced as shown on Fig. 8(b). Low current density leads to lower energy consumption, but in actual use, it is necessary to seek the balance between the energy saving and the $\mathrm{NO_3}^-$ treatment requirements to gain the highly appreciable treatment performance.

3.3.2. Effect of chlorine concentration

As can be seen in Fig. S1, the main product of the electrocatalytic reduction of NO₃⁻ was NH₄⁺, which is also a contaminant that needed to be removed. The removal of NH3 is primarily takes place through direct oxidation on the anode surface and indirect oxidation mediated by active chlorine in the solution bulk. In the direct oxidation process, the NH₃ is first adsorbed on the surface of the anode and subsequently oxidized to N₂, as shown in Eq. (8) [30]. However, NH₄⁺ cannot be oxidized directly at the anode. Hence it is necessary to adjust the solution pH to reach the pK_a of the NH₄⁺/NH₃ acid-base pair (i.e., pH > 9.25). [32] In addition to the direct oxidation of NH₃ on the anode, the generated active chlorine species (Eqs. (9) and (10)) at the anode could also be involved in the oxidative transformation of NH₄⁺ to nitrogen gas (Eq. (11)) or NO₃⁻ (Eq. (12)) as well as the oxidation of NO₂⁻ to NO₃⁻ (Eq. (13)) in the homogeneous solution, so-called indirect (or mediated) oxidation [30]. Furthermore, HOCl reacts also on NH₃ to form chloramines (Eqs. (14)-(16)). Compared with direct ammonia oxidation, the chlorine-mediated ammonia oxidation is much more efficient and feasible due to the abundant presence of chloride ion in wastewaters.

$$2NH_3 + 6OH^- \rightarrow N_2 + 6H_2O + 6e^-$$
 (8)

$$2Cl^- \rightarrow Cl_2 + 2e^- \tag{9}$$

$$Cl_2 + H_2O \rightarrow HOCl + H^+ + Cl^-$$
 (10)

$$2NH_4^+ + 3HOCl \rightarrow N_2 + 3H_2O + 3Cl^- + 5H^+$$
 (11)

$$NH_4^+ + 4HOCl \rightarrow NO_3^- + H_2O + 6H^+ + 4Cl^-$$
 (12)

$$NO_2^- + HClO \rightarrow NO_3^- + H_2O + H^+ + Cl^-$$
 (13)

(14)

$$HOCl + NH2Cl \rightarrow NHCl2 + H2O$$
 (15)

 $HOCl + NH_3 \rightarrow NH_2Cl + H_2O$

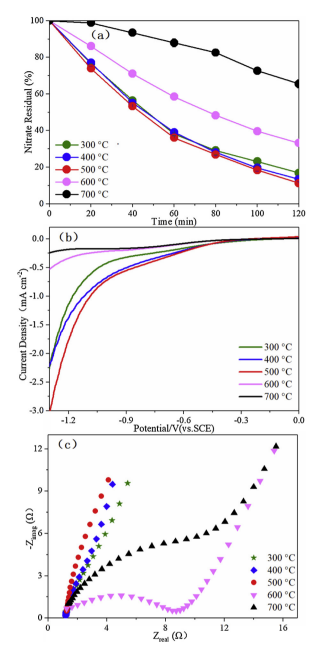


Fig. 6. NO_3^- concentration decay kinetics at different calcination temperature (a) under experimental conditions: $[NO_{3^-}]_0 = 50 \, \text{mg L}^{-1}$, pH = 7.0, current density = $10 \, \text{mA cm}^{-2}$), LSV curves at different calcination temperature (b) in $0.1 \, \text{M SO}_4^- + 50 \, \text{mg L}^{-1} \, NO_3^-$ solution at a scan rate of $100 \, \text{mV s}^{-1}$ and (c) EIS Nyquist plots of the cathodes at different calcination temperature by applying an impedance amplitude of $10 \, \text{mV}$ with the frequency range from 10^5 to 10^{-2} Hz.

$$HOCl + NHCl2 \rightarrow NCl3 + H2O$$
 (16)

Chloride ions are widely presented in drinking water and industrial water. For instance, the concentration of chloride ions in the bulk drug manufacturing industrial wastewater and zinc refinery wastewater reached 3119 and $10,000\,\mathrm{mg\,L}^{-1}$, respectively [35]. Considering the role of chloride ions in oxidation of NH₄+, various concentrations of chloride ions were added to the solution in order to study its effect. As shown in Fig. 9(a), the NO₃- removal efficiency was almost unaffected by the presence of chloride ions. But, as the concentration of chloride ions increased from 0 to 2000 mg L⁻¹, the formation of NH₄+ progressively decreased (Fig. 9(b)). For instance, when the concentration of chloride ions increased to 2000 mg L⁻¹, the generated NH₄+ ions ratio

initially slightly increased to 7% within 40 min but then gradually decreased to approximately 0 after 100 min reaction time. Besides, it appears from Fig. 9(c) that the formation of NO₂- could be deteriorated with increasing chloride ions concentration from 0 to 2000 mg $\rm L^{-1}$, with the highest NO₂- generation of 0.47%. It can also be seen from Fig. S6 that the TN removal efficiency and N₂ selectivity of Co₃O₄-TiO₂/Ti cathode increased with the increase of chloride ions concentration. When the chloride ions concentration reached 2000 mg $\rm L^{-1}$, the TN removal efficiency and N₂ selectivity reached 82% and 100%, respectively.

NH₄⁺ formation curves plotted in Fig. 9(d) clearly show that its amount formed at Co₃O₄/Ti cathode (49%) was about twice of that formed at Co₃O₄-TiO₂/Ti cathode (24%) after 120 min reaction at the same chloride ions concentration of $1000 \,\mathrm{mg}\,\mathrm{L}^{-1}$. Fig. S6 shows that the TN removal efficiency and N2 selectivity for Co3O4-TiO2/Ti cathode were 60% and 71%, respectively, while the corresponding values for Co₃O₄/Ti cathode were 42% and 46%, respectively. The difference in NT removal efficiency between Co₃O₄-TiO₂/Ti and Co₃O₄/Ti cathodes in the presence of chloride ions was much larger than in the chloridefree cases as observed in Fig. S1. The indirect oxidation mechanism of NH₄⁺ was similar to the intensively studied breakpoint chlorination [31]. First, the NH₄⁺ reacts with the active chlorine (HOCl or OCldepending to the solution the pH) to produce intermediate products such as monochloramine, dichloramine and tri-chloramine, and were finally oxidized to N2 [32]. It is worth noting that the oxidation process of the NH₄⁺ began to occur only when active chlorine/ammonia mole ratio reaches 1.5 [33]. Specially, Ding et al. [34] found that higher active chlorine/ammonia mole ratio (> 5) brought about 100% removal of NH₄⁺, while less efficient removal efficiencies (nearly 50%) were achieved with lower mole ratio (< 2). Thus, it can be inferred that electrochemical ammonia oxidation in these reaction systems was closelv related with the active chlorine/ammonia mole ratio. As demonstrated in Fig. 1, the efficiency of NO₃- reduction to NH₄ with Co₃O₄/ Ti cathode was higher than that with Co₃O₄-TiO₂/Ti cathode. This would result in less mole ratio of active chlorine/ammonia in the former case compared with that in the latter case. Therefore, as shown in Fig. 9(d), the indirect oxidation of ammonia was more favorable for the Co₃O₄-TiO₂/Ti cathode mediated electrochemical system.

3.4. The NO_3 removal mechanism

3.4.1. Direct reduction

Electrochemical reduction of NO₃ is a complex process involving a range of intermediates and broad valence changes (from +5 to -3). The process mainly includes direct reduction and indirect reduction. Direct reduction is mainly achieved by electrons and H+, and indirect reduction is mainly caused by atomic H* [35]. This mechanism was verified in the present electrochemical reaction system by CV tests as depicted in Fig. 10(a). When 50 mg L⁻¹ NO₃⁻ was added to the reaction solution, the peak current density of -2.4 and -4.0 mA cm⁻² appeared at the potential of -0.9 and -1.2 V/SCE, respectively, at TiO₂/Ti cathode. These reduction peaks can be attributed to the electron transfer from the cathode to NO3- because no peak was observed in the absence of NO₃⁻. Thus, it is rational that approximately 45% NO₃⁻ was reduced at TiO₂/Ti cathode (Fig. 1(a)). According to the previous studies [36], these two reduction peaks correspond to the reduction of NO₃⁻ to NO₂ and the formation of NH₄+, respectively. Furthermore, single reduction peak does not mean single electron transfer [37]. In contrast, the peaks were completely disappeared in the case of Co₃O₄-TiO₂/Ti cathode, regardless of the presence or absence of NO₃⁻. Notably, the experimental results presented in Fig. 1 show that the NO₃ reduction efficiency was greatly increased in the presence of Co₃O₄. Therefore, the disappearance of the reduction peak was not ascribed to the poor catalytic activity of the Co₃O₄ toward the electrocatalytic reduction of NO₃⁻. This peculiar phenomenon just became a breakthrough point to understand the mechanism of the NO3- electrocatalytic reduction by

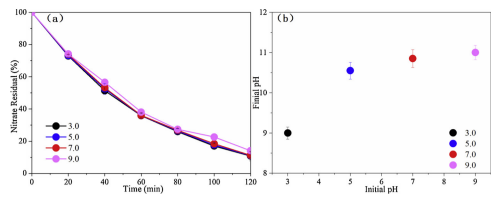


Fig. 7. NO_3^- concentration decay kinetics at different pH (a) and Final solution pH for various initial pH after 120 min reaction (b). Experimental conditions: $[NO_3^-]_0 = 50 \text{ mg L}^{-1}$, current density = 10 mA cm⁻².

Co₃O₄. It is known that two processes, i.e., heterogeneous charge transfer and diffusional mass transport, mainly determine the shape of the CV curve [38]. The CV curve can reflect the heterogeneous charge transfer from the electrode to electroactive species. However, it fails to reflect the charge transfer and valence change inside the electrode. Co₃O₄ contains Co²⁺ ions in tetrahedral interstices and Co³⁺ ions in octahedral interstices. Thus, it can be inferred that NO₃⁻ probably did not directly receive electrons from the cathode via the circuit, but from the "electron porter" of the structural Co²⁺ ions. This proposition can be validated by the density functional theory (DFT) calculation results, which can be seen in Fig. 11. Co₃O₄ (222) was selected considering XRD detection and NO₃ - reduction experiments. The DFT calculation results show that the O atom in NO₃ and the Co atom on the surface of Co₃O₄ (222) are thermodynamically favorably bonded with the required energy of -9.04 eV. Besides, the differential charge density distribution indicates that electrons can transfer from the Co atoms on the surface of Co₃O₄ (222) to NO₃⁻. Differential charge density analysis further showed that the O atoms in NO₃ obtained 0.41, 0.65, and 0.57 electrons, respectively, and the N atoms lost 0.82 electrons, and a total of 0.81 electrons were transferred from the Co atom to NO₃⁻.

In order to further confirm this, the electrochemical reduction of NO_3^- at Co_3O_4 -Ti O_2 /Ti cathode was investigated at different cathode potentials, and the experimental results are shown in Fig. 10(b). Nearly no NO_3^- was removed at the applied potential of -0.5 and -0.7 V/SCE within 120 min. The NO_3^- removal efficiency of 36% and 40% were attained at -1.0 and -1.2 V/SCE, respectively. Besides, NO_3^- removal efficiency finally raised to 98% after 120 min electrolysis when the applied potential was further decreased to -1.3 V/SCE. However, the NO_3^- reduction efficiency was decreased to approximately 94% when further decreasing the cathode potential to -1.5 V/SCE. This can be attributed to the enhancement of hydrogen evolution reaction at this potential and the hindrance of hydrogen bubbles to the electron transfer at the cathode surface [39]. The strong dependence of NO_3^-

reduction on the cathode potential corresponding to reduction peaks of CV curve is in agreement with the literature [40]. This confirms the above conjecture that electrons were transferred from the power source to the cathode surface through a circuit, in which $\mathrm{Co^{3}}^{+}$ was reduced to $\mathrm{Co^{2}}^{+}$ via accepting an electron. Then nitrogen-containing contaminants, such as $\mathrm{NO_{3}}^{-}$, receive the electrons from the structural $\mathrm{Co^{2}}^{+}$ ions. This particular reaction process cannot be directly detected by CV test, explaining the disappearance of the reduction peak in Fig. 10(a) and the strong voltage-dependent phenomenon in Fig. 10(b).

In the above process, the ratio $\mathrm{Co}^{2+}/\mathrm{Co}^{3+}$ would definitely change. To further confirm this statement, XPS analysis was used. The XPS wide-survey scan of Co(2p) is illustrated in Fig. 10(c). The peaks at 780 and 795 eV (binding energy) should be assigned to Co(2p3/2) and Co $(2p_{1/2})$, respectively [41]. The Co $(2p_{3/2})$ /Co $(2p_{1/2})$ ratio value of almost 2:1 was consistent with the previous literature report [42]. The peaks of Co²⁺ were located at 781.5 and 796.6 eV, while the peaks at 780 and 794.8 eV should be assigned to Co³⁺ [43,44]. Besides, the peaks at 786.8 and 804 eV were attributed to the shake-up satellite peaks of Co²⁺ [45]. Moreover, the Co²⁺/Co³⁺ ratios of the pristine and used Co₃O₄-TiO₂/Ti cathodes were calculated as 1.49 and 0.42, respectively, which indicated the transformation of Co²⁺ to Co³⁺ during the electrochemical NO₃ - reduction process. This confirms the electrochemical redox cycle Co²⁺-Co³⁺-Co²⁺ and the electron transfer from Co²⁺ to NO₃⁻ during the electrocatalytic NO₃⁻ reduction process.

3.4.2. Indirect reduction

In the indirect pathway, electrons from the cathode could produce surface-adsorbed atomic H^* via the reduction of protons, which is known as Volmer process [15]. Here, we employed CV technique to qualitatively detect the formed H^* during the electrochemical reaction process, which has been well introduced by Jiang et al. [46] and Liu et al. [47]. First, the cathode was repeatedly scanned for activation in a

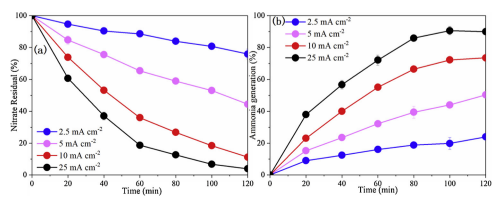


Fig. 8. NO₃ concentration decay kinetics (a) and NH₄ generation (b) at different current density. Experimental conditions: [NO₃-]₀ = 50 mg L⁻¹, pH = 7.0.

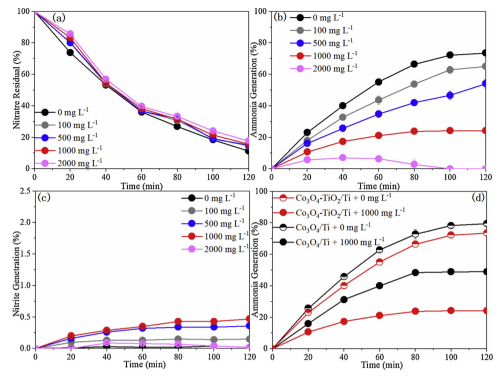


Fig. 9. Effect of Cl $^-$ concentration on NO $_3^-$ decay kinetics (a), NH $_4^+$ generation (b), NO $_2^-$ generation (c) with Co $_3$ O $_4$ -TiO $_2$ /Ti cathode, and NH $_4^+$ generation with Co $_3$ O $_4$ -TiO $_2$ /Ti and Co $_3$ O $_4$ /Ti cathodes (d). Experimental conditions: [NO $_3^-$] $_0^-$ = 50 mg L $^-$ 1, pH = 7.0, current density = 10 mA cm $^-$ 2.

potential range of -0.20 to 0.90 V/SCE in 0.1 M $\rm Na_2SO_4$ electrolyte at sweep rate of 100 mV s⁻¹. Electrochemical analysis did not start until a stable CV curve was obtained [46]. As shown in Fig. 10(d), the starting potential was set from -1.20 to -1.35 V/SCE and the termination potential was 0.45 V/SCE. As reported in the previous study involving C-Pd cathode for nitrate reduction [46]; the lower the starting potential, the stronger the corresponding oxidation peak. And the oxidation peaks of molecular hydrogen and atomic H* were located at -0.80 to -0.60 V/SCE and -0.10 to -0.00 V/SCE, respectively. Fig. 10(d)

shows that at different initial potentials, reduction peaks were observed at the potentials from -0.6 to $-0.1\,V/SCE$, corresponding to the oxidation of molecular H_2 and H^* , respectively. After adding 10 mM TBA, the intensity of the oxidation peak corresponding to atomic H^* immediately weakened. This markedly suppressed the availability of the formed H^* for the NO_{3^-} reduction.

Fig. 10(e) and (f) show LSV curves and NO_3^- reduction results, respectively, after the addition of $10\,\text{mM}$ TBA. As illustrated in Fig. 10(e), the current intensity was significantly enhanced after the

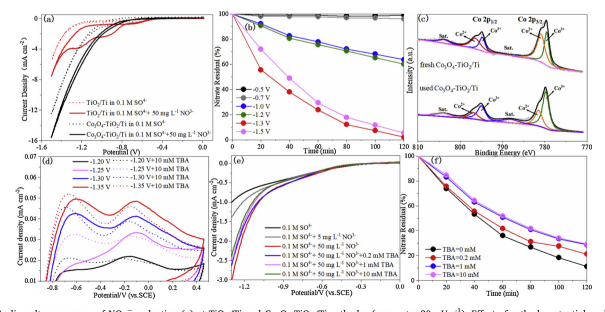


Fig. 10. Cyclic voltammograms of NO_3^- reduction (a) at TiO_2/Ti and Co_3O_4 - TiO_2/Ti cathodes (scan rate: 20 mV s^{-1}); Effect of cathode potential on NO_3^- concentration decay kinetics (b) at Co_3O_4 - TiO_2/Ti cathode (Experimental conditions: $[NO_3^-]_0 = 50 \text{ mg L}^{-1}$, pH = 7.0, current density $= 10 \text{ mA cm}^{-2}$); XPS spectra of Co 2p for fresh and used Co_3O_4 - TiO_2/Ti (c); Cyclic voltammograms obtained with Co_3O_4 - TiO_2/Ti cathode (d) in 0.1 M SO_4^- in the presence (dotted line) and absence (full line) of 10 mM TBA; LSV curves of different electrolyte (e); and concentration decay kinetics of NO_3^- at Co_3O_4 - TiO_2/Ti cathode (f) with various TBA concentrations (Experimental conditions: $[NO_3^-]_0 = 50 \text{ mg L}^{-1}$, pH = 7.0, current density $= 10 \text{ mA cm}^{-2}$).

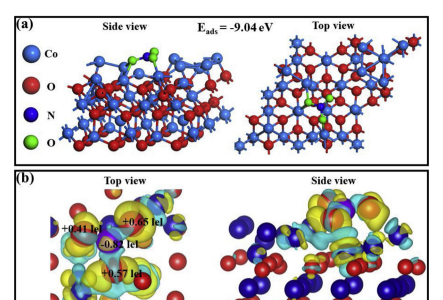


Fig. 11. (a) Adsorption configuration of NO_3^- on Co_3O_4 (222) surface and corresponding adsorption energy, (b) Differential charge density distribution of NO_3^- on Co_3O_4 (222) surface. Electron accumulation and electron deletion are represented by yellow and blue, respectively (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

addition of 5 and 50 mg $\rm L^1~NO_3^-$. However, the current intensity was decreased as TBA was added, which was more pronounced at higher TBA concentration. In addition, as shown in Fig. 10(f), increasing TBA concentration could gradually inhibit the $\rm NO_3^-$ removal efficiency. When TBA concentration was 0.2 mM, the $\rm NO_3^-$ removal efficiency was decreased by approximately 10%, and it was further decreased by 18% at TBA concentration of 1 mM. But negligible difference in the $\rm NO_3^-$ reduction efficiency between 1 and 10 mM TBA was observed, suggesting nearly complete hindering of the contribution of the formed $\rm H^*$ to $\rm NO_3^-$ reduction when TBA concentration was 1 mM. Therefore, we can reasonably speculate that approximately 18% $\rm NO_3^-$ reduction proceeded via the indirect pathway mediated by the atomic $\rm H^*$ during the electrocatalytic reduction of $\rm NO_3^-$ process.

Atomic H* would accordingly be generated first by Volmer mechanism (Eq. (17)), which can explain the increase of solution pH observed for the Co₃O₄-TiO₂-PVP cathode. In some noble metal catalysts (e.g., Pd and Ag)-based reaction systems, atomic H* would be favorably preserved for NO₃ reduction [12]. As a result, after the addition of 1 mM and 10 mM TBA in the Pd-mediated process, the NO₃ removal rate constant was reduced by 38% and 81%, respectively, which fully demonstrates the importance of atomic H* for NO₃⁻ reduction. For Pdbased catalysts, the high efficiency of the use of protons to generate atomic H* at low overpotentials (Eq. (17)) and the strong adsorption of atomic H* to Pd atoms make Eqs. (18) and (19) difficult to occur [47,48]. Even if atomic H* is converted to H₂, Pd atoms can cleave H₂ and regenerate atomic H* (this mechanism belongs to catalytic reduction) [12]. Through such electrocatalytic reduction processes, the indirect reduction of NO₃ by Pd-based catalysts is dominant. In contrast, in present Co₃O₄-based electrochemical system, the produced H* was inclined to transform to H2 by Heyrovsky and Tafel routes (Eqs. (17)–(19)) [15]. Consequently, as for Co₃O₄-TiO₂-PVP cathode, the indirect reduction mediated by atomic H* accounted for only 18% of NO₃ reduction. Thus, the role of atomic H* in the electrocatalytic reduction of NO₃⁻ at the Co₃O₄-TiO₂-PVP electrode was secondary, while the direct electron transfer was mainly responsible for NO₃ reduction. In view of this, future research should focus on improving the atomic H* adsorption capacity of non-precious metal catalysts.

$$H_2O + e^- \rightarrow H^* + OH^- \text{ (Volmer)}$$
 (17)

$$H_2O + H^* + e^- \rightarrow H_2(g) + OH^- \text{ (Heyrovsky)}$$
 (18)

$$2H^* \rightarrow H_2(g)$$
 (Tafel)

3.4.3. Proposed reaction pathway

It was observed from Fig. 1 that the $\mathrm{NH_4}^+$ concentration gradually increased over the reaction time for all cathodes containing $\mathrm{Co_3O_4}$ catalyst as it was the main product of electrocatalytic $\mathrm{NO_3}^-$ reduction, while $\mathrm{NO_2}^-$ was the minor intermediate product at a relatively low concentration. According to the aforementioned discussion and the experimental results, there are two main pathways (i.e., Co-mediated direct reduction and atomic H*-mediated indirect reduction) responsible for the electrocatalytic reduction of $\mathrm{NO_3}^-$ by the $\mathrm{Co_3O_4}$ bearing cathodes investigated in this study.

It is known that the reduction of NO_3^- to NO_2^- (Eq. (20)) is the kinetically rate-determining step for the electrocatalytic reduction of NO_3^- . This step actually consists of the reactions set presented through Eqs. (21)–(23) which is known as an electrochemical-chemical-electrochemical path [35]. The elementary electron transfer reactions lead to the generation of $NO_{3\,(ad)}^2$ and $NO_{2\,(ad)}^2$, which are adsorbed short-lived intermediates. Then, $NO_{2\,(ad)}^2$ converts to an unstable dianion radical, namely $NO_{2\,(ad)}^2$, by direct electron transfer (Eq. (24)), which would quickly hydrolyzes to form $NO_{(ad)}$ (Eq. (25)) [35]:

$$NO_3^- + 2H^+ + 2e^- \rightarrow NO_2^- + H_2O E^0 = 0.01 \text{ V/SHE}$$
 (20)

$$NO_{3(ad)}^- + e^- \rightarrow NO_{3(ad)}^{2-} E^0 = -0.89 \text{ V/SHE}$$
 (21)

$$NO_{3(ad)}^{2-} + H_2O \rightarrow NO_{2(ad)} + 2OH - k = 5.5 \times 10^4 \text{ s}^{-1}$$
 (22)

$$NO_{2'(ad)} + e^- \rightarrow NO_{2(ad)}^- + H_2O E^0 = 1.04 \text{ V/SHE}$$
 (23)

$$NO_{2(ad)}^{-} + e^{-} \rightarrow NO_{2(ad)}^{2-} E^{0} = -0.47 \text{ V/SHE}$$
 (24)

$$NO_{2(ad)}^{2-} + H_2O \rightarrow NO_{(ad)} + 2OH$$
 (25)

According to the previous studies, the stepwise reduction of $NO_{(ad)}$ eventually forms NH_4^+ , which is regarded as an electrochemical-electrochemical path, as depicted in Eqs. (26)–(31) [49]:

$$NO_{(ad)} + H^{+} + e^{-} = HNO_{(ad)} E^{o} = -0.78 \text{ V/SHE}$$
 (26)

$$HNO_{(ad)} + H^{+} + e^{-} \rightarrow H_{2}NO_{(ad)} E^{o} = 0.52 \text{ V/SHE}$$
 (27)

$$H_2NO_{(ad)} + H^+ + e^- \rightarrow H_2NOH_{(ad)} E^0 = 0.90 \text{ V/SHE}$$
 (28)

$$H_2NOH + H^+ = H_3NOH^+$$
 (29)

$$H_2NOH + 2H^+ + 2e^- \rightarrow NH_3 + H_2O E^0 = 0.42 \text{ V/SHE}$$
 (30)

(19)

$$NH_3 + H^+ = NH_4^+ pK_a = 9.25$$
 (31)

The direct electron transfers in above reactions are mostly single electron transfer processes. The above experimental results demonstrated that the redox process ${\rm Co^{2+}-Co^{3+}-Co^{2+}}$ accelerates the electron transfer during ${\rm NO_3}^-$ reduction. The source of electrons for ${\rm NO_3}^-$ reduction is directly derived from ${\rm Co^{2+}}$ instead of the cathode through the circuit. ${\rm Co^{2+}}$ gives an electron to ${\rm NO_3}^-$ or its daughter intermediates with the formation of ${\rm Co^{3+}}$, and then ${\rm Co^{3+}}$ would transform to ${\rm Co^{2+}}$ via obtaining an electron from the cathode surface. Thus, during the whole reaction, ${\rm Co_3O_4}$ mainly acts as an electron shuttle rather than a reducing agent. Moreover, ${\rm H^+}$ is consumed in many ${\rm NO_3}^-$ reduction steps, which is consistent with the pH variation as shown in Fig. 7(b).

At the same time, due to the strong reducibility of atomic hydrogen $(E_{'(H+/H)} = -2.31 \text{ V/SHE})$ [49], the indirect reduction path of NO_3^- can not be ignored. Atomic H* involved in Eqs. (32)–(37).

$$NO_{3(ad)}^{-} + 2H^* \rightarrow NO_{2(ad)}^{-} + H_2O$$
 (32)

$$NO_{2(ad)}^{-} + H^* \rightarrow NO_{(ad)} + OH^{-}$$
 (33)

$$NO_{(ad)} + 2H^* \rightarrow N_{(ad)} + H_2O$$
 (34)

$$N_{(ad)} + H^* \rightarrow NH_{(ad)} \tag{35}$$

$$NH_{(ad)} + H^* \rightarrow NH_{2(ad)}$$
 (36)

$$NH_{2(ad)} + H^* \rightarrow NH_{3(ad)} \tag{37}$$

4. Conclusions

A novel Co₃O₄-TiO₂/Ti cathode was synthesized for removal of NO₃ from water by its electrocatalytic reduction of. It was proved that adding a certain amount of PVP in the coating liquid could improve the dispersibility of the catalyst and therefore promote the electrocatalytic reduction of NO₃⁻. More importantly, the release of toxic Co ions into the solution was almost completely prohibited during the NO₃- reduction process due to the stabilization effect of TiO2. The introduction of a certain amount of chloride ions into the electrolyte significantly increased the N₂ selectivity and total nitrogen removal efficiency. When the concentration of chloride ions reached 2000 mg L-1, almost no NH₄⁺ was detected after 100 min of electrolysis. The electrocatalytic mechanism of NO3 - reduction consisted of direct reduction and indirect reduction, which are mediated by the Co²⁺-Co³⁺-Co²⁺ redox cycle and atomic H*, respectively. Notably, the former route played a superior role in NO₃⁻ reduction over the latter one. Generally, considering the abundant chloride ions in industrial/municipal wastewaters, the developed Co₃O₄-TiO₂/Ti cathode in this study might be a promising alternative for the electrocatalytic treatment of aqueous effluents loaded with NO3-.

Acknowledgments

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Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.apcatb.2019.05.016.

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